

# MetaboAnalystを用いた PCA, PLS-DA

<https://www.metaboanalyst.ca/>

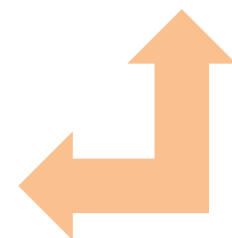
# データファイルの作り方

各サンプルが各行になるようにする。行と列を入れ替えることも可能

1列目にID、2列目にサンプルの分類情報、3列目以降に変数データを記入する

id	class	Q1	Q2	Q3	Q4	Q5	Q6	Q7	Q8
1	M	100	100	50	70	10	100	100	
2	F	80	100	90	100	0	90	100	
3	F	50	100	100	100	0	70	80	
4	F	100	100	50	50	0	70	100	
5	F	94	80	98	30	60	98	99	
6	F	99	100	80	70	0	10	60	
7	M	50	40	80	50	0	80	40	
8	F	55	50	69	100	0	0	15	
9	F	25	0	100	100	0	25	75	
10	F	30	75	80	60	0	90	80	
11	M	85	100	85	90	0	100	40	
12	M	23	74	87	72	0	68	82	
13	F	70	100	80	100	10	60	60	
14	F	55	90	100	80	30	50	42	
15	F	50	80	100	80	0	30	100	
16	F	90	50	70	70	0	0	40	
17	M	80	30	100	100	2	30	95	

転置可能



# データファイルの作り方

カンマ区切りファイル(csv)にしたいデータだけを記載した、エクセルシートを作る。

そのシートを閲覧している状態で、「名前を付けて保存」から、カンマ区切り(csv)を選んで保存する。

# MetaboAnalystへアクセス

click here to startをクリック



**MetaboAnalyst 5.0** - user-friendly, streamlined metabolomics data analysis

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## News & Updates

- Check out our [MetaboAnalyst 5.0 paper](#) now available on the 2021 NAR web server issue; **NEW**
- For multi-omics analysis, try our [OmicsAnalyst](#) for various data-driven approaches, or [OmicsNet](#) for knowledge-based integration; **NEW**
- Improved size calculation for heatmap visualization in Statistics module (08/05/2021); **NEW**
- Added support for VIP plot for Orthogonal Partial Least-Squares (OPLS) in Statistics module (07/15/2021); **NEW**
- Minor bug fixes based on user feedback (05/31/2021); **NEW**
- Improved functional meta-analysis data upload page for more flexible data processing (04/27/2021);
- Fixed the issue with synchronized 3D scatter plots for scores and loadings (04/12/2021);
- Upgraded to PrimeFaces 10 and fixed the issue for multi-group figure legend (04/06/2021);
- Added support for merging technical replicates with different mathematical models and QC measures (02/18/2021);

[Read more...](#)

**Click here to start**

## Overview

MetaboAnalyst is a comprehensive platform dedicated for metabolomics data analysis via user-friendly, web-based interface. Over the past decade, MetaboAnalyst has evolved to become the most widely used platform (>300,000 users) in the



# Statistical Analysisをクリック

## Module Overview

Input Data Type	Available Modules (click on a module to proceed, or scroll down for more details)					
Raw Spectra (mzML, mzXML or mzData)				LC-MS Spectra Processing		
MS Peaks (peak list or intensity table)			Functional Analysis	Functional Meta-analysis		
Annotated Features (compound list or table)		Enrichment Analysis	Pathway Analysis	Joint-Pathway Analysis	Network Analysis	
Generic Format (.csv or .txt table files)	Statistical Analysis	Biomarker Analysis	Time-series/Two-factor Analysis	Statistical Meta-analysis	Power Analysis	Other Utilities

どちらでもOK

➤ [Statistical Analysis](#)

This module is only used statistical and machine learning methods including t-tests, ANOVA, PCA, PLS-DA and Orthogonal PLS-DA. It also provides clustering and visualization tools to create dendrograms and heatmaps as well as to classify data based on random forests and SVM.

➤ [Spectral Analysis](#)

This module allows users to upload raw LC-MS spectra (mzML, mzXML or mzData) to be processed using our optimized workflow based on MetaboAnalystR - OptiLCMS. The module supports common LC-MS platforms. The result peak intensity table can be used for statistical and functional analysis.

➤ [Biomarker Analysis](#)

This module performs various biomarker analyses based on receiver operating characteristic (ROC) curves for a single or multiple biomarkers using well-established methods. It also allows users to manually specify biomarker models and perform new sample prediction.

➤ [Functional Analysis \(MS Peaks\)](#)

This module accepts high-resolution LC-MS spectral peak data to perform metabolic pathway enrichment analysis and visual exploration based on the well-established mummichog algorithm. It currently supports 26 organisms including Human, Mouse, Zebrafish, *C. elegans*, and other species.

➤ [Pathway Analysis \(targeted\)](#)

This module supports pathway analysis (integrating enrichment analysis and pathway topology analysis) and visualization for 26 model organisms, including Human, Mouse, Rat, Cow, Chicken, Zebrafish, *Arabidopsis thaliana*, Rice, Drosophila, Malaria, *S. cerevisiae*, *E.coli*, and others species.

➤ [Functional Meta-analysis \(MS peaks\)](#)

This module aims to identify robust **functional profiles** across multiple global metabolomics datasets via two approaches: 1) integrating functional profiles from independent studies conducted under compatible LC-MS conditions; or 2) pooling peaks from complementary instruments within the same studies.

➤ [Time-series/Two-factor Analysis](#)

➤ [Enrichment Analysis](#)

➤ [Joint Pathway Analysis](#)

# 準備したCSVファイルをアップロード

必要に応じて、Formatなどを選択する

Submitボタンを押す

## 1) Upload your data

Tab-delimited text (.txt) or comma-separated values (.csv) file:

Data Type: ☒ Concentrations ☐ Spectral bins ☐ Peak intensity table

Format:

Data File:  okonomi.csv

**Submit**

# データチェックが行われる 問題なければskipで先に進む

## Data Integrity Check:

1. Checking the class labels - at least three replicates are required in each class.
2. If the samples are paired, the pair labels must conform to the specified format.
3. The data (except class labels) must not contain non-numeric values.
4. The presence of missing values or features with constant values (i.e. all zeros).

### Data processing information:

Checking data content ...passed.

Samples are in rows and features in columns

The uploaded file is in comma separated values (.csv) format.

The uploaded data file contains 33 (samples) by 29 (compounds) data matrix.

Samples are not paired.

2 groups were detected in samples.

Only English letters, numbers, underscore, hyphen and forward slash (/) are allowed.

Other special characters or punctuations (if any) will be stripped off.

All data values are numeric.

A total of 0 (0%) missing values were detected.

By default, missing values will be replaced by 1/5 of min positive values of their corresponding variables

Click the **Skip** button if you accept the default practice;

Or click the **Missing value imputation** to use other methods.

Edit Groups

Missing Values

→ Proceed

# 必要に応じて、データの標準化(normalization)、変換(transformation)、スケーリング方法を選択する

## Normalizeボタンを押す

### Normalization overview:

The normalization procedures are grouped into three categories. The sample normalization allows general-purpose adjustment for differences among your sample; data transformation and scaling are two different approaches to make individual features more comparable. You can use one or combine them to achieve better results.

#### Sample Normalization

☒ None

☐ Sample-specific normalization (i.e. weight, volume) [Specify](#)

☐ Normalization by sum

☐ Normalization by median

☐ Normalization by reference sample (PQN) [Specify](#)

☐ Normalization by a pooled sample from group [Specify](#)

☐ Normalization by reference feature [Specify](#)

☐ Quantile normalization

#### Data transformation

☒ None

☐ Log transformation (generalized logarithm transformation or glog)

☐ Cube root transformation (takes the cube root of data values)

#### Data scaling

☒ None

☐ Mean centering (mean-centered only)

☐ Auto scaling (mean-centered and divided by the standard deviation of each variable)

☐ Pareto scaling (mean-centered and divided by the square root of the standard deviation of each variable)

☐ Range scaling (mean-centered and divided by the range of each variable)

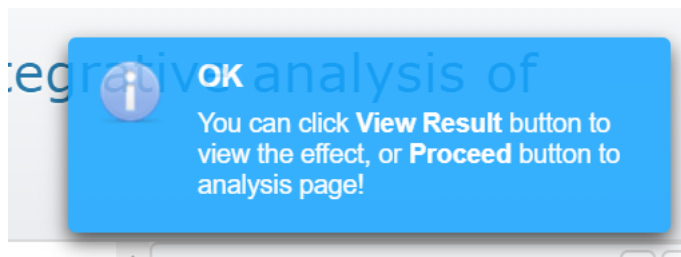
Normalize

View Result

Proceed



# 右上にこのウィンドウが出ればOK



# アクティブになったProceedボタンを押す

**Data scaling**

☒ None

☐ Mean centering (mean-centered only)

☐ Auto scaling (mean-centered and divided by the standard deviation of each variable)

☐ Pareto scaling (mean-centered and divided by the square root of the standard deviation of each variable)

☐ Range scaling (mean-centered and divided by the range of each variable)

Normalize View Result **Proceed**

# 主成分分析 Principal Component Analysis (PCA) をクリック

Select an analysis path to explore :

## Univariate Analysis

[Fold Change Analysis](#) [T-tests](#) [Volcano plot](#)

One-way Analysis of Variance (ANOVA)

[Correlation Heatmaps](#) [Pattern Search](#) [Correlation Networks \(DSPC\)](#)

## Chemometrics Analysis

[Principal Component Analysis \(PCA\)](#)

[Partial Least Squares - Discriminant Analysis \(PLS-DA\)](#)

[Sparse Partial Least Squares - Discriminant Analysis \(sPLS-DA\)](#)

[Orthogonal Partial Least Squares - Discriminant Analysis \(orthoPLS-DA\)](#)

## Feature Identification

[Significance Analysis of Microarray \(and Metabolites\) \(SAM\)](#)

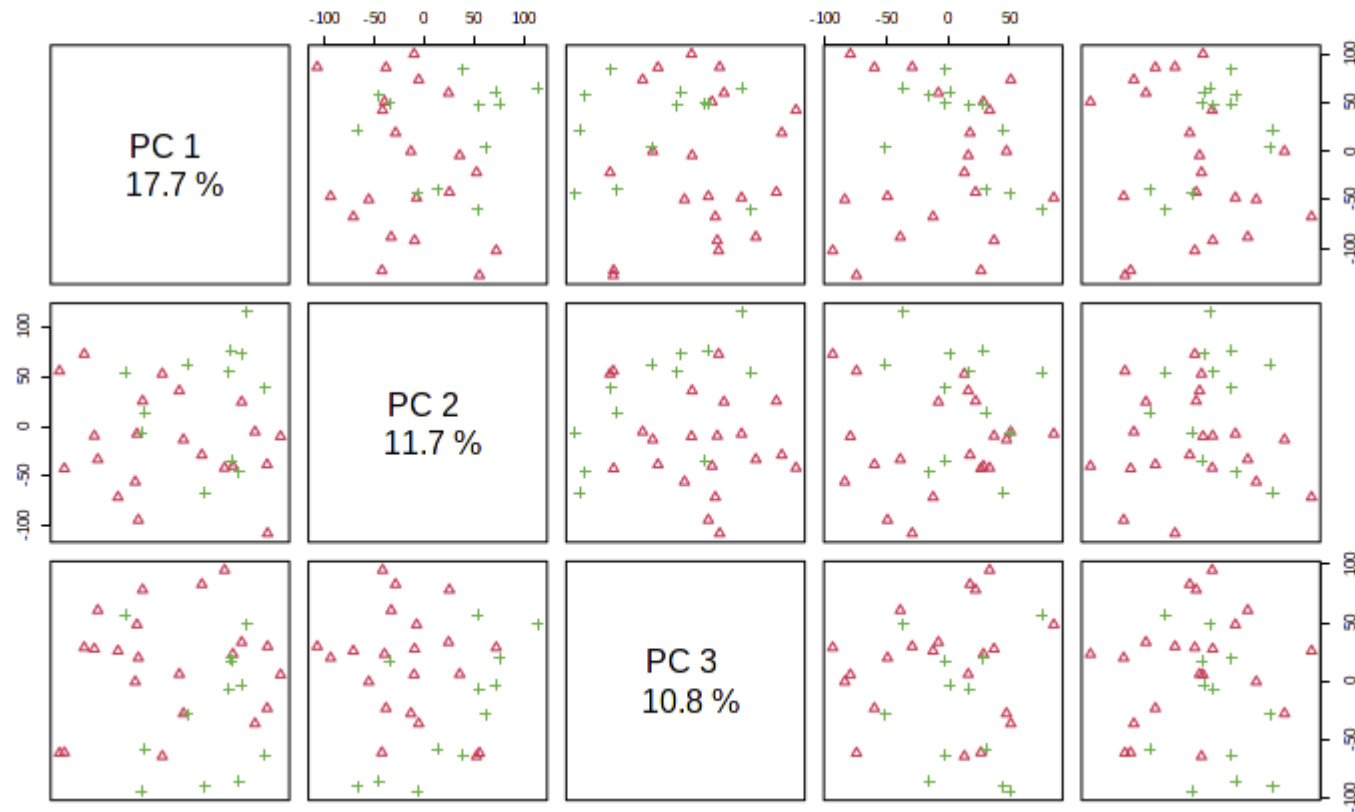
[Empirical Bayesian Analysis of Microarray \(and Metabolites\) \(EBAM\)](#)

## Cluster Analysis

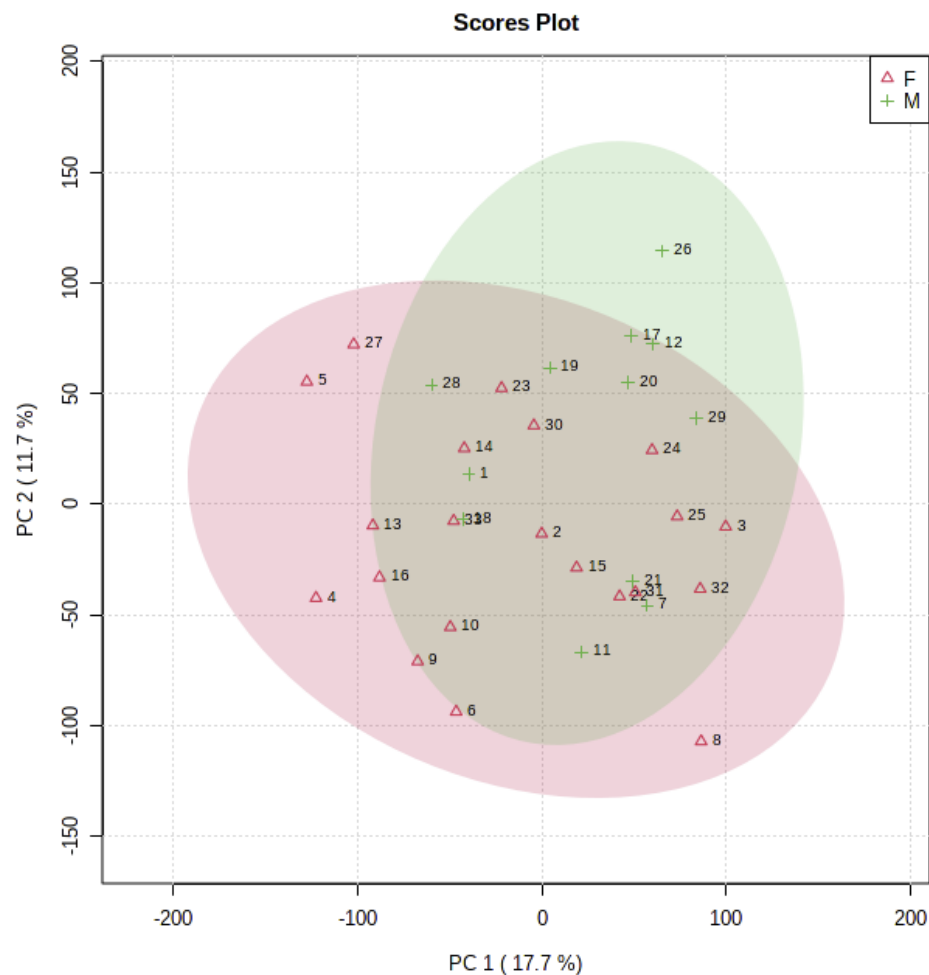
Hierarchical Clustering: [Dendrogram](#) [Heatmaps](#)

# 結果画面が表示される

## Principal Component Analysis (PCA)

[Overview](#)[Scree Plot](#)[2D Scores Plot](#)[Loadings Plot](#)[Synchronized 3D Plots](#)[Biplot](#)Display pairwise score plot for top  PCs

# 2D Scores Plot



Specify PC on X-axis:

Specify PC on Y-axis:

Display 95% confidence regions:

☒

Display sample names:

☒

Use grey-scale colors:

☐

Flip Image

☐

X axis

☒

Y axis

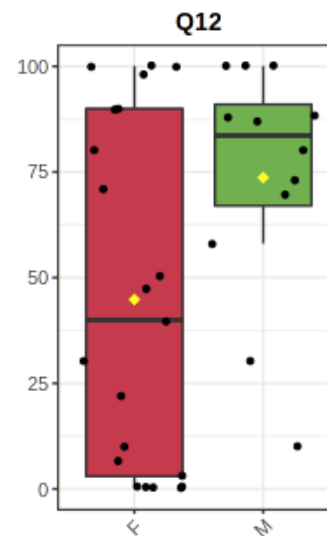
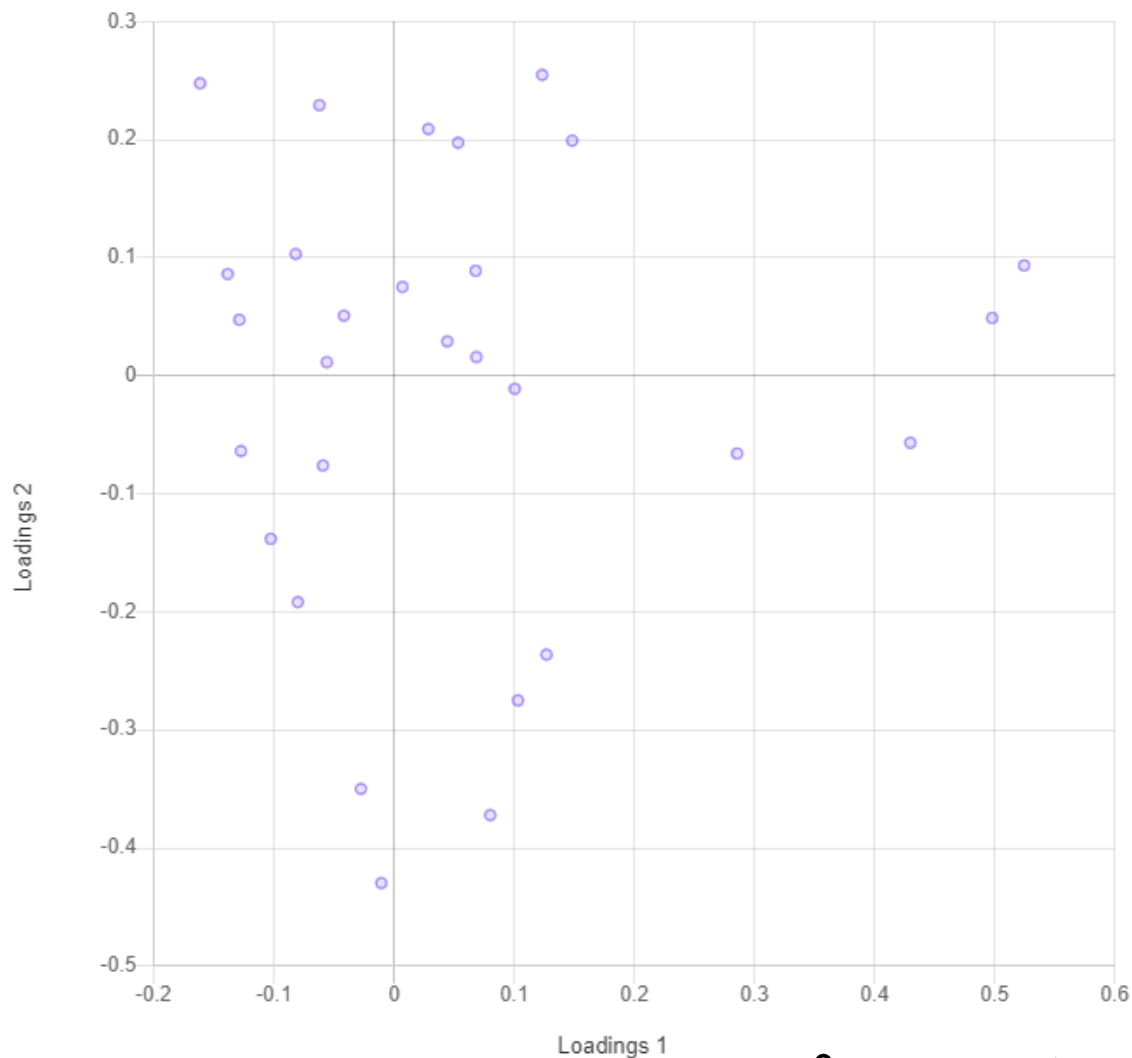
☐

All

Display sample  
nameにチェックが入っ  
ていると、サンプル名が  
一緒に表示される。

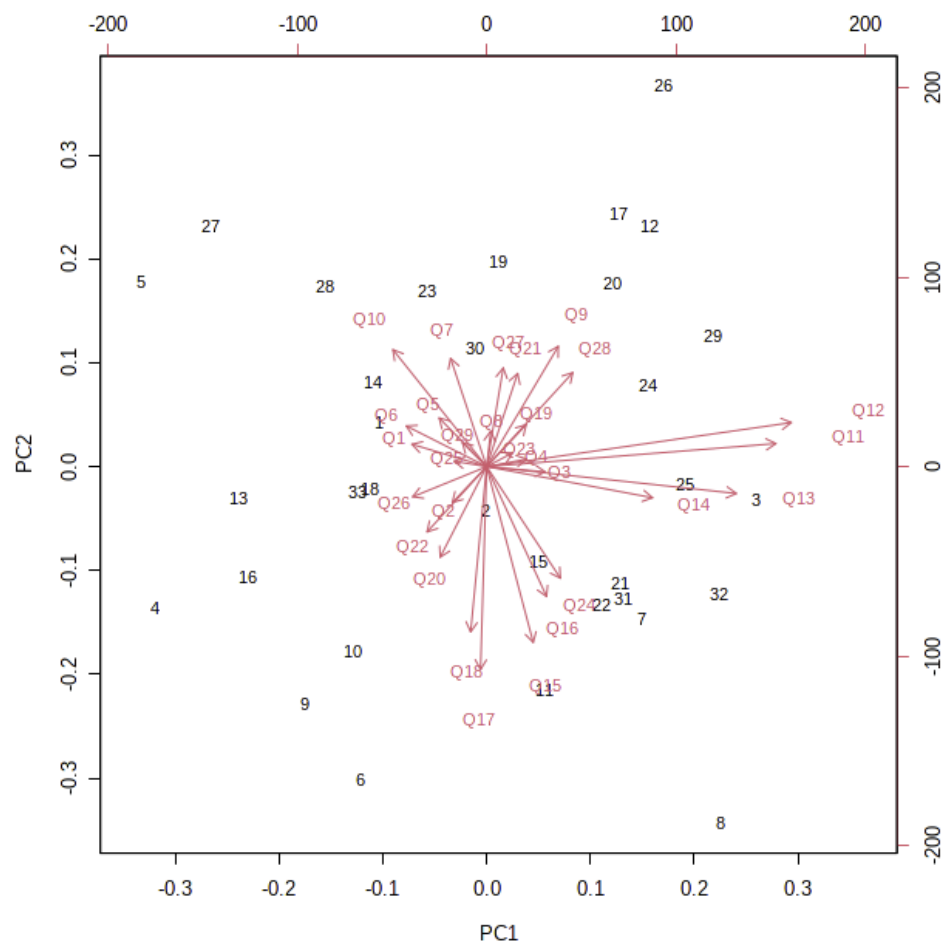
※条件を変えた後に「Update」ボタン  
を押して反映させます。

# Loadings Plot



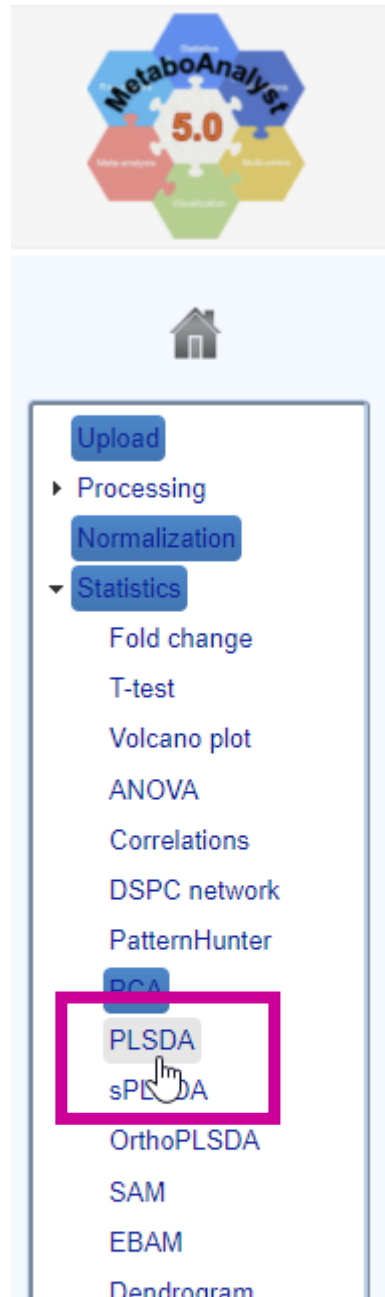
プロットをクリックすると、その変数の特徴が表示される。

# Biplot

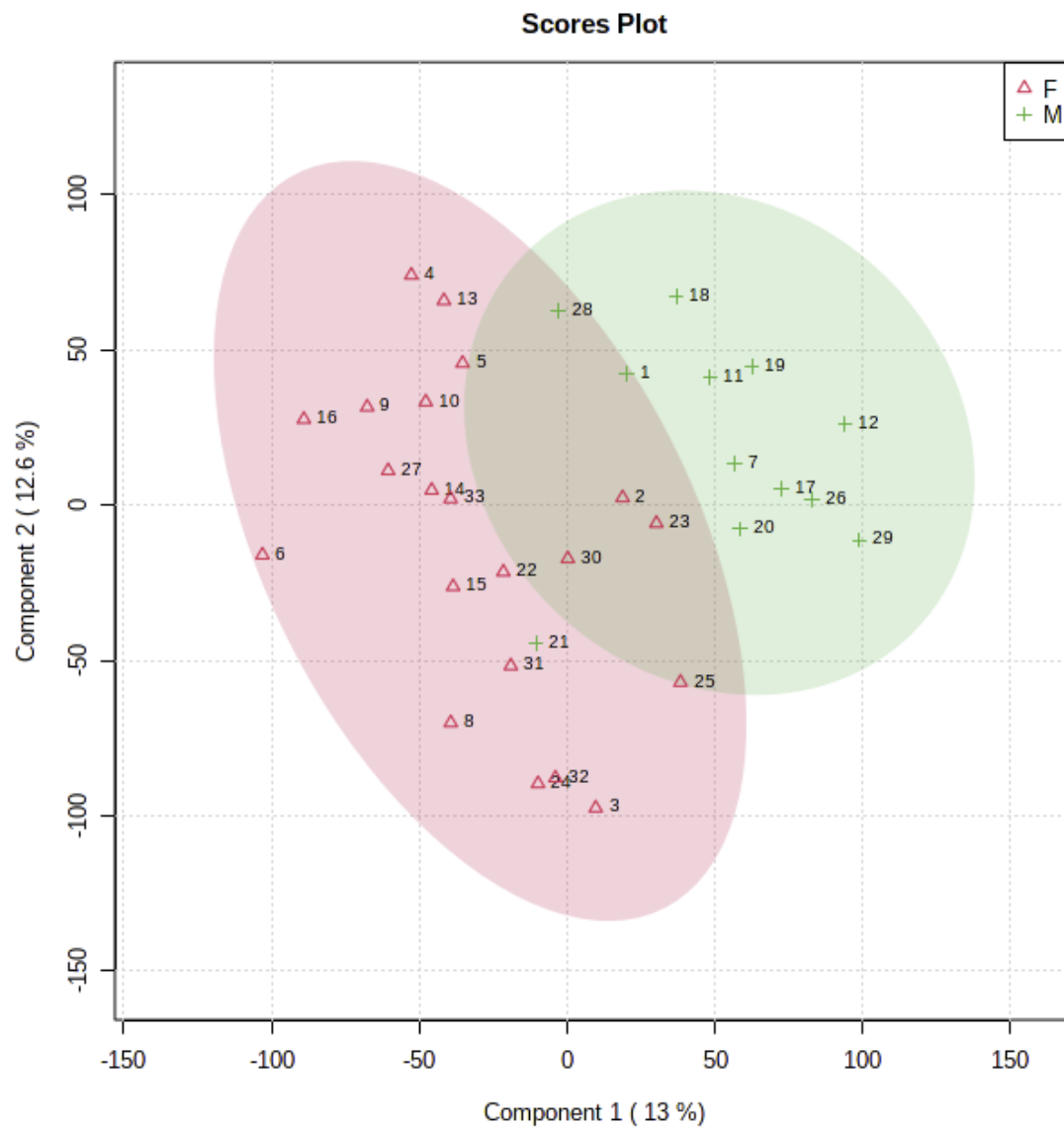


スコアプロットとローディングプロットが一緒に描かれたもの。

# PLSDA

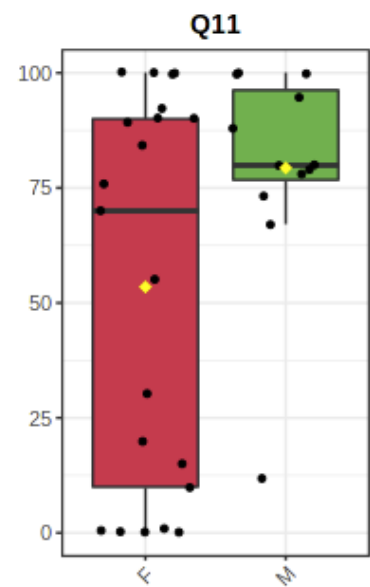
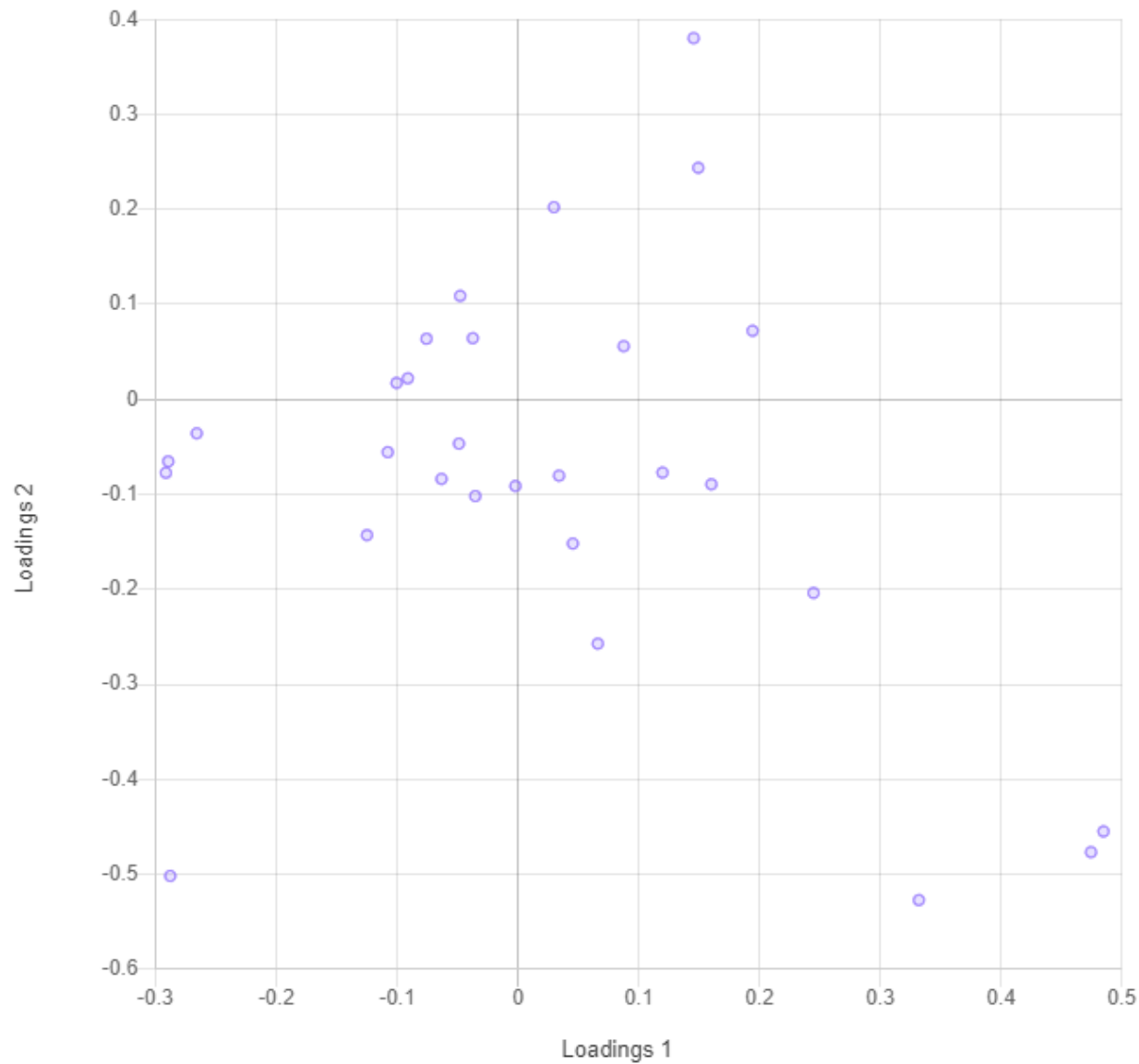


# 2D Scores Plot

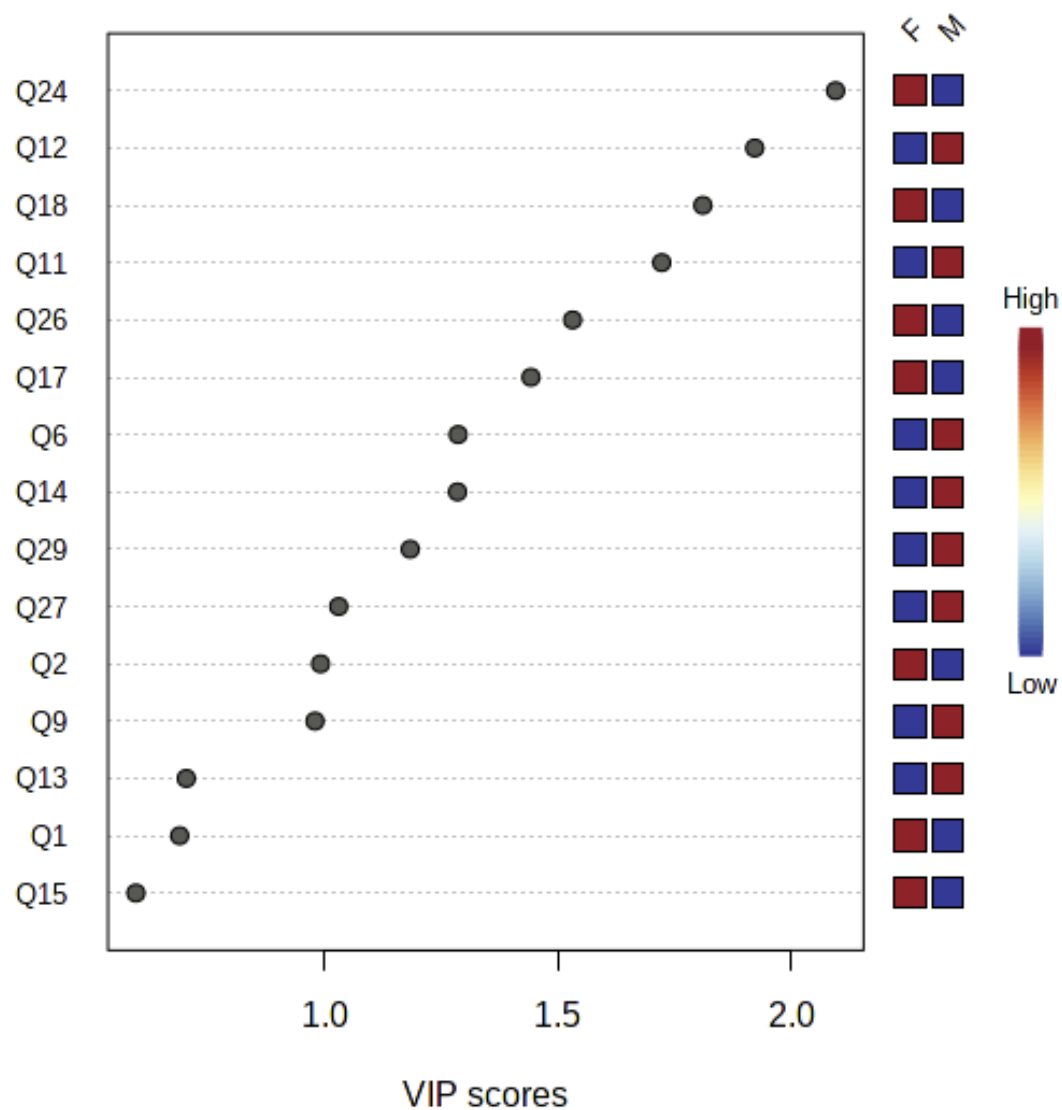




# Loadings Plot



# VIP值



# 累積寄与率と構築したモデルの精度(Q)

